REMARKS

Rejection of the claims under 35 U.S.C.§103

The Examiner maintains the rejection of the claims under 35 U.S.C.§103 as being obvious over Dianov et al. combined with McCarthy et al. US '176. Applicant traverse this rejection and withdrawal thereof if respectfully requested.

Applicants contend that a skilled artisan is not motivated to combine the references for the following reasons.

A. There is no motivation to combine the references because the proposed modification would make Dianov *et al.* unsatisfactory for its intended purpose.

The Dianov et al. reference teaches synthetic oligonucleotides, which each contain a single base that serves as a glycosylase substrate. This modified base is introduced into the oligonucleotides during artificial synthesis. The Examiner asserts that the oligonucleotides of Dianov et al. may be modified according to the teaching of the '176 reference. That is, the single aberrant base of Dianov et al. may be introduced into the oligonucleotides via enzymatic extension, and then subsequently, a 3'-OH end of an extendable DNA fragment may be formed.

However, applying the modification of the '176 reference to the oligonucleotides of Dianov et al. will result in more than a single base modification. According to the '176 reference at column 5, lines 49-53 "amplification will typically involve amplifying a target nucleic acid sample using a combination of normal DNA precursor nucleotides and one or more modified precursor nucleotide(s) where the modified precursor nucleotide replaces one of the normal precursor nucleotides." For example, as shown in Figure 1 of the '176 reference, dUTP replaces dTTP during amplification, and all of the thymine bases are replaced with uracil. Likewise, all of the thymine or guanine or cytosine or adenine bases in the Dianov et al. oligonucleotides would be replaced with the modified base of choice according to the '176 reference. As is evident from the Dianov et al. oligonucleotides depicted in Figure 2, (page 1607 of Dianov et al.), the enzymatic extension described in the '176 reference would result in multiple modified bases in the Dianov et al. oligonucleotides.

The incorporation of multiple modified bases into the oligonucleotides of Dianov et al. would render them unsatisfactory for their purpose. The purpose of the oligonucleotides in the

Application No. 09/673,739
Amendment dated August 21, 2006

After Final Office Action of December 23, 2005

Dianov et al. reference is to use them to demonstrate the mechanism of DNA repair of a single aberrant base. Because mechanisms of DNA repair vary and depend, inter alia, upon the number and location of aberrant bases that require repair, modifying the oligonucleotides of Dianov et al. according to the enzymatic extension procedure of the '176 reference, would not have allowed Dianov et al. to describe the pathway of single base repair.

More specifically, the purpose of the Dianov et al. reference is to answer the "critical question" (page 1611, first column second paragraph, Dianov et al.) of whether or not repair of a single aberrant base is 1 or 2 nucleotides in size. Dianov et al. answered this critical question by using synthetic double stranded oligonucleotides with sequences suited for restriction enzyme analysis. For example, in order to determine whether or not repair occurs 3' to a single modified base, Dianov et al. constructed a particular oligonucleotide sequence, termed "substrate c." This oligonucleotide is AT rich in the region of the single modified base. The oligonucleotide was designed in this manner so that if repair of the modified base occurred 3' to the AP site, a marker in a 13 nucleotide fragment would be apparent after SacI digestion.

Hence, modifying the Dianov *et al.* oligonucleotides according to the '176 reference would not only have resulted in numerous aberrant bases, thus rendering the purpose of Dianov *et al.* impossible, but in the case of one of the oligonucleotides, substrate c, the AT rich regions would have been lost, resulting in the loss of the *SacI* restriction site. These changes would have resulted in non-interpretable data.

Thus, the oligonucleotides of Dianov *et al.* cannot be modified by the enzymatic extension of the '176 reference without rendering the prior art invention unsatisfactory for its intended purpose. "If proposed modification would render the prior art invention being modified unsatisfactory for its intended purpose, then there is no suggestion or motivation to make the proposed modification." *In re Gordon*, 733 F.2d 900, 221 USPQ 1125 (Fed. Cir. 1984). Therefore, combining the Dianov *et al.* reference with the '176 reference is improper.

3 MAA/

Docket No.: 1377-0156P

Docket No.: 1377-0156P

B. There is no motivation to combine the references because the prior art does not suggest the desirability of the claimed invention.

Moreover, even if the Examiner were to find that the '176 modification of the Dianov et al. reference is not unsatisfactory for its intended purpose, the prior art must suggest the desirability of the claimed invention. "In determining the propriety of the Patent Office case for obviousness in the first instance, it is necessary to ascertain whether or not the reference teachings would appear to be sufficient for one of ordinary skill in the relevant art having the reference before him to make the proposed...combination..." In re Linter, 458 F.2d 1013, 1016, 173 USPQ 560, 562 (CCPA 1972). Applicants contend that the desirability of the instant invention is not suggested by the combination of the Dianov et al. reference and the proposed modification of the '176 reference.

The instant invention concerns the characterization of a nucleic acid molecule template of interest via the extension of a 3'-OH end of an extendable DNA fragment on the nucleic acid molecule template of interest. The present invention is used, inter alia, to identify one or more bases in a nucleic acid of interest that were previously unknown. Additionally, the extendable fragment may be used to determine whether or not a particular sequence contains a single nucleotide polymorphism or other mutation, which is unknown for a particular sample. The combination of the Dianov et al. reference with the modification of the '176 reference does not suggest these desirable features.

The Dianov et al. reference is concerned with DNA repair of a single aberrant base. "[t]he main finding in the present work...is that the prevalent repair reaction involves the filling-in of a gap of only a single nucleotide" (page 1610, column 2, paragraph 2). Additionally, the constitution and location of the single repaired base of Dianov et al., unlike the characterization of one or more bases in the nucleic acid template of interest of the present invention, is a base, which is known to the skilled artisan before extension of the alleged extendable fragment. That is, during the generation of the alleged extendable fragment of Dianov et al., as combined with the '175 reference, a known base is replaced with a modified base. Thus, the constitution of the replaced/repaired base is known because, according to the '176 reference, modified bases replace known bases (e.g. uracil replaces thymine).

4 MAA/

Application No. 09/673,739 Amendment dated August 21, 2006 After Final Office Action of December 23, 2005

Additionally, the location of the replaced/repaired base, also, is known before extension of the alleged extendable fragment of Dianov *et al*. The modified base of Dianov *et al*., which corresponds to the replaced base, is removed *via* glycosylase. Cleavage then occurs to form the alleged 3'-OH end of an extendable DNA fragment. Thus, the skilled artisan recognizes that the location of the replaced/repaired base is found on the strand complementary to the alleged 3'-OH end of an extendable DNA fragment, *i.e.* one base 3' to the end of the alleged 3'-OH end of the extendable DNA fragment on the opposite strand.

Subsequently, the alleged 3'-OH end of the extendable DNA fragment is extended, but only to repair the previously identified base; no additional unknown bases are identified. ("These data demonstrate that the majority of DNA repair replication events in the *E. coli* cell extract involved the replacement of only one nucleotide." (page 1609, first column, lines 6-8. Dianov et al.)).

Furthermore, the Dianov et al. reference suggests that no further bases could be identified with the alleged 3'-OH end of the extendable DNA fragment using the template as shown in Figure 9. The pathway for single base repair, which is depicted on the left side of Figure 9 (the right side depicts the pathway that rarely occurs) shows that the alleged 3'-OH end of the extendable DNA fragment could not be further extended because it is blocked by a second 5' to 3' fragment.

Therefore, the skilled artisan is not provided with combined references that suggest that unknown bases can be identified with the alleged 3'-OH end of the extendable DNA fragment of Dianov *et al.* Applicants contend that an inventive step would be required to determine how the alleged 3'-OH end of the extendable DNA fragment of Dianov *et al.*, as modified by the '176 reference, could be used to identify unknown bases on selected templates.

Thus, because the desirable features of the present invention, that of identifying unknown bases, is not suggested by the Dianov *et al.* reference using the proposed modification of the '176 reference, the combination of the references is improper and the 103(a) rejection should be withdrawn.

5 MAA/

Docket No.: 1377-0156P

Application No. 09/673,739 Amendment dated August 21, 2006

After Final Office Action of December 23, 2005

In view of the above remarks, Applicants believe the pending application is in

condition for allowance.

Should there be any outstanding matters that need to be resolved in the present

application, the Examiner is respectfully requested to contact MaryAnne Armstrong, PhD Reg.

No. 40,069at the telephone number of the undersigned below, to conduct an interview in an

effort to expedite prosecution in connection with the present application.

If necessary, the Commissioner is hereby authorized in this, concurrent, and future

replies to charge payment or credit any overpayment to Deposit Account No. 02-2448 for any

additional fees required under 37.C.F.R. §§1.16 or 1.14; particularly, extension of time fees.

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Respectfully submitted,

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Docket No.: 1377-0156P

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6 MAA/